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30	Under the Paperwork Reduction Act of 1995, no.	nersons	are required to respond to a collection			PTO/SB/21 (02-04) 07/31/2006. OMB 0651-0031 ARTMENT OF COMMERCE a valid OMB control number.
	Utilier the Paperwork Reduction Act of 1995, No.	oersons	Application Number	10/767,813		
_ 1	TRANSMITTAL		Filing Date	January 29, 2004		
CRAI	FORM		First Named Inventor	Zhang et al.		
	-		Art Unit	unassigned		
	(to be used for all correspondence after initial filing	g)	Examiner Name	unassigned		
	29		Attorney Docket Number	AHPWA1DUSA		
	Total Number of Pages in This Submission 29				1	
		ENC	LOSURES (Check all tha	t apply)		
	Fee Transmittal Form Fee Attached Amendment/Reply After Final Affidavits/declaration(s) Extension of Time Request Express Abandonment Request Information Disclosure Statement Certified Copy of Priority Document(s) Response to Missing Parts/ Incomplete Application Response to Missing Parts under 37 CFR 1.52 or 1.53		Drawing(s) Licensing-related Papers Petition Petition to Convert to a Provisional Application Power of Attorney, Revocation Change of Correspondence Add Terminal Disclaimer Request for Refund CD, Number of CD(s)	to Ted Appe of Ap Appe (Appe Propi Statu Othe Ident	chnology al Comments are al Comments Notice rietary In	ure(s) (please
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	I hereby certify that this correspondence is be sufficient postage as first class mail in an envithe date shown below.	eing fac elope a	simile transmitted to the USPTO addressed to: Commissioner for F	or deposited with the Patents, P.O. Box 1450	United S 0, Alexa	ndria, VA 22313-1450 on
	Typed or printed name					
	Signature				Date	

This collection of information is required by 37 CFR 1.5. The information is required to obtain or retain a benefit by the public which is to file (and by the USPTO to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.14. This collection is estimated to 2 hours to complete, including pathering, preparing, and submitting the completed application form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450.



AHPWA1DUSA

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Appln. No.

: 10/767,813

Confirmation No.: unassigned

Applicant

: Zhang et al.

Filed

: January 29, 2004

TC/A.U.

: unassigned

Examiner

: unassigned

Customer No.

: 38199

Title

: CYCLOCARBAMATE DERIVATIVES AS PROGESTERONE

RECEPTOR MODULATORS

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INFORMATION DISCLOSURE STATEMENT

Sir:

Applicants submit to the Examiner the attached Form PTO/SB/08A/B document listing and this paper pursuant to 37 CFR § 1.56 and § 1.97-1.98. Form PTO/SB/08A/B is attached and copies of documents GH-GK, HB-HE, HK-IG, DM, and EO are enclosed herewith. This Information Disclosure Statement is being submitted more than three months from the filing date of this application but before the receipt of a first Office Action.

The Director is hereby authorized to charge any deficiency in any fees due with the filing of this paper or credit any overpayment in any fees to our Deposit Account Number 08-3040.

Express Mail No. EU531570788US

REMARKS

Listed below are previously filed, and co-pending, US Patent applications. These applications and the present application are commonly owned:

- (1) U.S. Patent Application No. 09/552,633, filed April 19, 2002, now U.S. Patent No. 6,509,334, issued January 21, 2003 (item GC)
 - U.S. Patent Application No. 09/948,309, filed September 6, 2001, now U.S. Patent No. 6,566,358 issued May 20, 2003 (item GF)
 - U. S. Patent Application No. 10/386,799, filed March 12, 2003, now U.S. Patent No. 6,713,478, issued March 30, 2004 (item GK)
- (2) U.S. Patent Application No. 09/552,632, filed April 19, 2000, now U.S. Patent No. 6,391,907, issued May 21, 2002 (item FB)
 - U.S. Patent Application No. 10/014,173, filed December 11, 2001, now U.S. Patent No. 6,608,068, issued August 18, 2003 (item GI)
 - U.S. Patent Application No. 10/456,892, filed June 6, 2003 (U.S. Patent Publication No. US-2003-0220388-A1, published November 27, 2003 (item HD))
- (3) U.S Patent Application No. 09/552,352, filed April 19, 2000, now U.S. Patent No. 6,417,214, issued July 9, 2002 (item FE)
 - U.S. Patent Application No. 10/131,379, filed April 24, 2002 (U.S. Patent Publication No. US-2003-0008909-A1, published January 9, 2003 (item HH))
- (4) U.S. Patent Application No. 09/552,033, filed April 19, 2000, now U.S. Patent No. 6,355,648, issued March 12, 2002 (item EG)
 - U.S. Patent Application No. 10/022,467, filed October 30, 2001, now U.S. Patent No. 6,521,657, issued February 18, 2003 (item GD)
 - U.S. Patent Application No. 10/117,156, filed April 5, 2002 (U.S. Patent Publication No. US-2002-0169198-A1, published November 14, 2002 (item HB))
 - U.S. Patent Application No. 10/253,380, filed September 24, 2002, now U.S. Patent No. 6,583,145, issued June 24, 2003 (item GG)

- U.S. Patent Application No. 10/420,276, filed April 22, 2003 (U.S. Patent Publication No. US-2003-0225109-A1, published December 12, 2003 (item HE))
- (5) U.S. Patent Application No. 09/552,354, filed April 19, 2000, now U.S. Patent No. 6,436,929, issued August 20, 2002 (item FG)
 - U.S. Patent Application No. 10/140,034, filed May 6, 2002 (U.S. Patent Publication No. U.S.-2003-0092711-A1, published May 15, 2003 (item HJ))
- (6) U.S. Patent Application No. 09/552,546, filed April 19, 2000, now U.S. Patent No. 6,380,235, issued April 30, 2002 (item FA)
 - U.S. Patent Application No. 10/074,768, filed February 12, 2002 (U.S. Patent Publication No. US-2002-0115853-A1, published August 22, 2002 (HF))
- (7) U.S. Patent Application No. 09/552,630, filed April 19, 2000, now U.S. Patent No. 6,339,098, issued January 15, 2002 (item EF)
- (8) U.S. Patent Application No. 09/552,036, filed April 19, 2000, now U.S. Patent No. 6,306,851, issued October 23, 2001 (item EC)
 - U.S. Patent Application No. 09/906,875, filed July 17, 2001, now U.S. Patent No. 6,441,019, issued August 27, 2002 (item FH)
- (9) U.S. Patent Application No. 09/552,356, filed April 19, 2000, now U.S. Patent No. 6,369,056, issued April 9, 2002 (item EJ)
 - U.S. Patent Application No. 10/050,287, filed January 16, 2002 (U.S. Patent Publication No. US-2002-0111355-A1, published August 15, 2002 (item IB))
- (10) U.S. Patent Application No. 09/552,629, filed April 19, 2000, now U.S. Patent No. 6,358,948, issued March 19, 2002 (item EI)
 - U.S. Patent Application No. 10/023,063, filed December 17, 2001, now U.S. Patent No. 6,693,103, issued February 17, 2004 (item GJ)
- (11) U. S. Patent Application No. 09/552,544, filed April 19, 2000, now U.S. Patent No. 6,407,101, issued June 18, 2002 (item FD)

- U.S. Patent Application No. 10/043,513, filed January 9, 2002, now U.S. Patent No. 6,562,857, issued May 13, 2003 (item GH)
- U.S. Patent Application No. 10/342,719, filed January 15, 2003 (U.S. Patent Publication No. US-2003-0158182-A1, published August 21, 2003 (item HC))
- (12) U.S. Patent Application No. 09/552,357, filed April 19, 2000, now U.S. Patent No. 6,498,154, issued December 24, 2002 (item GA)
- (13) U.S. Patent Application No. 09/552,037, filed April 19, 2000, now U.S. Patent No. 6,399,593, issued June 4, 2002 (item FC)
- U.S. Patent Application No. 09/552,350, filed April 19, 2000, now U.S.Patent No. 6,444,668, issued September 3, 2002 (item FJ)
 - U.S. Patent Application No. 10/141,792, filed May 9, 2002 (U.S. Patent Publication No. US-2003-0045511-A1, published March 6, 2003 (item HI))
- (15) U.S. Patent Application No. 09/552,631, filed April 19, 2000, now U.S. Patent No. 6,329,416, issued December 11, 2001 (item EE)
 - U.S. Patent Application No. 09/977,790, filed October 15, 2001, now U.S. Patent No. 6,503,939, issued January 7, 2003 (item GB)
- U.S. Patent Application No. 09/552,355, filed April 19, 2000, now U.S.
 Patent No. 6,423,699, issued July 23, 2002 (item FF)
 - U.S. Patent Application No. 10/091,222, filed March 1, 2002 (U.S. Patent Publication No. US-2002-0151531-A1, published October 17, 2002 (item HG))
- U.S. Patent Application No. 09/552,545, filed April 19, 2000, now U.S.
 Patent No. 6,380,178, issued April 30, 2002 (item EK)
- (18) U.S. Patent Application No. 09/552,358, filed April 19, 2000, now U.S. Patent No. 6,462,032, issued October 8, 2002 (item FK)
 - U.S. Patent Application No. 10/153,393 filed May 22, 2002, now U.S. Patent 6,544,970 issued April 8, 2003 (item GE)
- (19) U.S. Patent Application No. 09/552,038, filed April 19, 2000, now U.S. Patent No. 6,319,912, issued November 20, 2001 (item ED)

- U.S. Patent Application No. 09/552,353 filed, April 19, 2000, now U.S.
 Patent No. 6,358,947, issued March 19, 2002 (item EH)
- U.S. Patent Application No. 10/601,442, filed June 23, 2003
 (U.S. Patent Publication No. US-2004-0006122-A1, published January 8, 2004 (item IC))
- U.S. Patent Application No. 10/601,968, filed June 23, 2003
 (U.S. Patent Publication No. US-2004-0014798-A1, published January 22, 2004 (item ID))
- U.S. Patent Application No. 10/601,438, filed June 23, 2003
 (U.S. Patent Publication No. US-2004-0002535-A1, published January 1, 2004 (item IE))
- U.S. Patent Application No. 10/601,481, filed June 23, 2003
 (U.S. Patent Publication No. US-2004-0006060-A1, published January 8, 2004 (item IF))

Applicants have also provided comments on the documents cited in this Information Disclosure Statement that were published in a language other than English. Specifically, Applicants have provided the following comments for the following documents.

- (1) English translations of documents EQ and AL were not available. However, document BJ, U.S. Patent No. 5,414,088, issued May 9, 1995, corresponds to document EQ, International Patent Publication No. WO 91/04974, published April 18, 1991. Similarly, document BB, i.e. U.S. Patent No. 4,831,027, issued May 16, 1989, corresponds to document AL, German Patent No. 3,633,861, issued April 7, 1988. Further, document ES, is the corresponding Chemical Abstracts entry to document BB. Brief remarks on these documents follow:
 - (EQ) This document describes the preparation of bicyclobenzimidazoles and their use to inhibit erythrocyte and thrombocyte aggregation. Thus, these compounds are used to treat conditions such as arterial occlusive or ischaemic conditions, venous insufficiency or diabetes mellitus.

- (AL) This document describes the preparation of imidazobenzoxazinones, and pharmaceutical compositions containing them. These compounds are assertedly useful in the treatment of cardiovascular effects, particularly cardiotonic activity and antithrombotic activity, with little effect on blood pressure.
- (2) English translations of documents CN, BM, BL, and DM were not available. Abstracts of these documents were obtained and were provided as documents BW, AV, AS, and DU respectively. Brief remarks on these documents follow:
 - (CN) This document describes the preparation and use of imidazopyridine derivatives as platelet agglutination inhibitors, anti-allergic, anti-inflammatory sedative, cardiac and cardiovascular vasodilators.
 - (BM) This document describes the preparation of pharmaceutical compositions of derivatives of benzimidazoles and azabenzimidazoles, which assertedly have cardiotonic, vasodilating, anti-hypertensive, anti-aggregation, and anti-ulcer activity. These compositions are assertedly useful as cardiotonics, vasodilators, anti-hypertensives, and anti-platelet aggregations agents. Further, these compounds assertedly have anti-ulcer activity and can be used in the treatment of gastroduodenal ulcers.
 - (BL) This document describes the preparation, manufacture, and use of heterocyclic, substituted azoles and azines as herbicides. Particular attention is drawn to their use for selective control of mono- and dicotyledonous weeds and are better tolerated than known compounds of similar structure.
 - (DM) This document refers to 2,3-dihydro-pyrido[1,2,3-de]-1,4-benzoxazine chloride compounds and a condensed hetercyclic ring structure which are assertedly useful as organic luminophores for marking agents.
 - English translations of documents GP, FP, and FS were not available. English language abstracts were provided on the front page of the documents. Brief remarks on these documents follow:

- (GP) This document describes the combination of one compound having progesterone-antagonistic properties with one compound having anti-estrogen properties. Each compound is used in a dose that would not inhibit ovulation by itself, but is effective when combined. The medicaments are discussed as being useful in female contraception.
- (FP) WO 95/11013 describes the use of at least one compound with progesterone antagonistic action and at least one compound with anti-estrogen action, together with partial agonistic action, for drugs assertedly useful in hormone replacement therapy.
- (FS) Vernin describes the preparation of 6-aryl- and 6-heteroaryl-2-ethylbenzothiazoles. The electrophilic character of the 2-ethyl-6-benzothiazolyl intermediate radicals is also discussed. A number of the 6-aryl and 6-heteroaryl-2-ethylbenzothiazoles were subsequently utilized to prepare corresponding quaternary salts and spriopyrans.
- (4) Abstracts for documents AM and AN were obtained and were provided as documents, BS and BR, respectively. Applicants have also enclosed an English language translation of AN and have provided brief remarks on these documents as follow:
 - (AM) DE 4,330,234 relates to female fertility control comprising intermittent administration of competitive progesterone antagonist together with daily of continuous administration of gestagen. The gestagen may be levonorgestrel, gestodene, desogestrel or cyproterone acetate and the antagonist may be mifepristone or onapriston.
 - (AN) DE 4,344,463 relates to a contraceptive pack comprising a combination of individual dosage units of competitive progesterone antagonist in an amount which does not inhibit ovulation or promote abortion and individual dosage units of a gestagen for sequential oral administration. Administration can be oral, topical or local.

- (5) An English translation of document HN was not available. However, document DK, US Patent No. 6,077,840, issued June 20, 2000, corresponds to document HN, International Patent Publication No. WO 98/27059 and is a parallel document to European Patent No. 947507 (document CL). A brief remark on this document follows:
 - (HN) This document describes a progesterone receptor binding inhibitor which is a tetrahydrobenzindolone derivative of the formula provided. The compound is described as useful as a carcinostatic agent.
- (6) English language abstracts of FL and GQ were provided on the front page of the documents. Applicants also provide English language translations of these documents. Brief remarks on these documents follow:
 - (FL) This document describes the preparation and use of new bicycloimidazoles. The compounds of the invention are particularly directed toward the use in drugs to prevent clumping of both erythrocytes and thrombocytes.
 - (GQ) This document describes the preparation and use of heterocyclically-substituted 1-indole carboxamides, their pharmaceutically acceptable salts, and use as cyclooxygenase-2 inhibitors.
- (7) English translations of documents CP, CQ, and DL were not available. However, document HK, US Patent No. 4,518,597, issued May 21, 1985, corresponds to document CP. Document EO, International Patent Publication No. WO-95/04048, published February 9, 1995, corresponds to document CQ. Document IA, US Patent No. 6,013,647, issued January 11, 2000, corresponds to document DL. Brief remarks on these documents follow:
 - (CP) This document describes novel benzoxazin-2-ones that have pharmacological properties in antithrombotic activity, which can be prepared using the methods for analogous compounds.

- (CQ) This document describes new benzoxazinedione derivatives useful with antibiotic agents to improve antibacterial effects against relatively resistant bacterial strains.
- (DL) This document describes the preparation and use of benzoxazinones and benzothiazinones, their pharmaceutically acceptable salts, and use as therapeutically active substances.
- (8) English translations of documents DN, DP, and DQ were not available. However, document BK, U.S. Patent No. 5,447,928, issued September 5, 1995, corresponds to document DN. Document CH, U.S. Patent No. 5,659,046, issued August 19, 1997, corresponds to document DP. Document FQ, International Patent Application No. WO95/20389, published August 3, 1995, corresponds to document DQ. Brief remarks on these documents follow:
 - (DN) This document describes optically pure or mixtures of benzoxazine compounds useful as neuronal calcium antagonists. These compounds are assertedly useful in treating damage following ischaemic attack, cardiac or respiratory arrest, cerebral thrombosis or embolism, cerebral senility, dementias, Alzheimer's disease, Huntington's chorea, soliv-ponto-cerebellar atrophy, amyotrophic lateral sclerosis, cranial and spinal trauma, preventing neuronal damage following convulsions, and in treating cancers, neurological changes to the AIDS, and diabetic retinopathies.
 - (DP) This document describes methods of preparing 2-perfluoroalkyl-3 oxazolin-5-one, which are assertedly effective against insects, mites, and nematodes.
 - (DQ) This document describes benzoxainone compounds of the general formula:

$$X \xrightarrow{X^1} R$$

where X=halogen

These compounds are assertedly useful for inhibiting the activity of HIV reverse transcriptase, prophylaxis, and treatment of HIV-infections and AIDS.

- (9) An English translation of document DO could not be obtained. However, this document corresponds to document CE, U.S. Patent No. 5,552,412, issued September 3, 1996; document EB, U.S Patent No. 6,204,286, issued March 20, 2001 (which is a continuation of document CE); document EA, U.S. Patent No. 6,153,622, issued November 28, 2000 (which is a continuation-in-part of document EB); document FI, U.S. Patent No. 6,441,193, issued August 27, 2002 (which is a continuation of document EB); and document HA, U.S. Patent Publication No. US-2001-0025051A1, published September 27, 2001 (which is a continuation of document FI). Brief remarks on this document follow:
 - (DO) This document describes compounds of the following structure:

These compounds, their optic and geometric isomers and nontoxic pharmaceutically acceptable acid-additive salts assertedly have estrogen-antagonistic/agonistic activity.

(10) English translations of documents CO and EL-EN could not be obtained.

However, English language abstracts of documents CO and EM-EN were provided as documents BT, BY, and CX. Further, documents BI, CF, and BG are the corresponding US patents for documents CO and EM-EN. The following comments for documents CO and EM-EN are also provided.

(CO) This document discusses compounds having the following structure:

These compounds are assertedly useful as having anti-arrhythmic activity, for anti-aggregation of blood-platelets, having beta-blocking properties, and for treating cardiac insufficiency, arrhythmia, migraines, and angina pectoris.

(EM) This document discusses compounds having the following structure:

$$(H_2C)$$
 (H_2C)
 $($

wherein:

X is H, F, Cl, Br, C_{1-6} alkyl, C_{3-8} cycloalkyl, NO_2 , CF_3 , CN, SH, $S(O)_mR^3$, OR^4 , COR^4 , or $CONR^4R^5$; and Y=H, F, Cl, Br, C_{1-6} alkyl, C_{3-8} cycloalkyl, NO_2 , CF_3 , CN, SH, $S(O)_qR^{17}$, OR^{18} , or $CONR^{18}R^{19}$.

These compounds are assertedly useful as analgesics, antiinflammatories, and antiarthritic agents, and to inhibit IL-1 biosynthesis (osteoporosis, periodontal disease, and tissue scarring) and immune dysfunctions (allergies and psoriasis). (EN) This document discusses compounds having the following structure:

$$R^3$$
 R^4
 R^5
 R^6
 R^2
 R^1
 R^6

wherein:

R = H, OH, or F; R^1 - $R^4 = H$, $C_{1^{-4}}$ alkyl, X, CF₃, phenyl, p-methylphenyl, or CF₃-phenyl; or $R^1 + R^2$, $R^2 + R^3$, or $R^3 + R^4$ are joined to from a benzo-fused ring.

These compounds are assertedly useful in the treatment of disorders responsive to the opening of potassium channels such as ischemia, convulsions, asthma, and traumatic brain injuries.

- (11) Applicants could not obtain an English translation or an English language abstract of document EL. However, from information provided on the Russian Patent Office web site Applicants were able to obtain an English language abstract of the priority application, i.e., Great Britain Patent Application No. 9326136. Further, document CG is the corresponding US patent to the priority application. The following comments for document EL are also provided.
 - (EL) This document discusses the following compounds:

$$R_3$$
 R_4
 R_5
 $R_1O)_n$
 R_2
 R_4
 R_5

 R_4 = H, OH, C_{1-6} alkoxy, C_{2-6} alkanoyloxy, carboxy, nitro, or NHR; and R_5 = H, C_{1-6} alkyl, or halogen.

These compounds are assertedly useful in inhibiting angiogenesis.

The month of publication for documents AR, AY, CU, CW, CY, DR, DV, DX, ES, ET, EU, EV, and EZ could not be determined. The years of publication for these documents are sufficiently earlier than the effective US filing date and/or the foreign priority date of the present application so that the particular month of publication of the documents is not an issue (MPEP 609).

The Examiner is respectfully requested to consider the enclosed documents identified in this paper and in the attached Form PTO/SB/08A/B during the course of examination of this application.

Respectfully submitted,

HOWSON AND HOWSON Attorneys for Applicants

Cathy A Kodroff

Registration No. 33,980

Spring House Corporate Center

Box 457

Spring House, PA 19477 Telephone: (215) 540-9200 Telefacsimile: (215) 540-5818

PTO/SB/08A (08-03)

Approved for use through 07/31/2006. OMB 0651-0031

U.S. Patent and Trademark Office; U.S. DEPARTMENT OF COMMERCE the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it contains a valid OMB control number.

Substitute for form 1449/PTO

Application Number 10/767,813 Filing Date January 29, 2004 First Named Inventor Zhang et al. Art Unit unassigned

INFORMATION DISCLOSURE STATEMENT BY APPLICANT

(Use as many sheets as necessary)

Examiner Name unassigned

Sheet 1 of 15 Attorney Docket Number AHPWA1DUSA

			U. S. PATENT	DOCUMENTS	
Examiner Initials*	Cite No.1	Document Number	Publication Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevant Passages or Relevant Figures Appear
	↓	Number-Kind Code ^{2 (# known)}			1 iguics / tppca/
	AA	^{US-} 3,526,621	11-01-1970	Bernardi	
-	AB	^{US-} 3,635,941	01-18-1972	Weaver	
	AC	^{US-} 3,635,964	01-18-1972	Skorcz	
	AD	^{US-} 3,917,592	11-04-1975	Kobzina	
,	AE	^{US-} 4,093,730	06-06-1978	Butti	
	AF	^{US-} 4,440,785	04-03-1984	Walsh	
	AG	^{US-} 4,617,302	10-14-1986	Robertson	
	АН	^{US-} 4,666,913	05-19-1987	Kuhla	
	AI	^{US-} 4,670,566	06-02-1987	Walsh	
	AJ	^{US-} 4,721,721	12-26-1988	Kuhla	
	AK	^{US-} 4,792,561	12-20-1988	Walker	
	ВА	^{US-} 4,822,794	04-18-1989	Spada	
	ВВ	^{US-} 4,831,027	05-16-1989	Narr	
	ВС	us- 4,853,473	08-01-1989	Fischer	
	BD	us- 4,933,336	06-12-1990	Martin	
	BE	us- 5,007,952	04-16-1991	Kume	
	BF	^{US-} 5,171,851	12-15-1992	Kim	
	BG	^{US-} 5,182,282	01-26-1993	Clemence	
	ВН	^{US-} 5,246,989	09-21-1993	Iwamoto ·	

		FORE	IGN PATENT DOCL	MENTS		
Examiner Initials*	Cite No.1		Publication Date	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevant Passages	١.
		Country Code ³ "Number ⁴ "Kind Code ⁵ (if known)	MM-DD-YYYY		Or Relevant Figures Appear	T ⁶
	AL	DE-3,633,861	04-07-1988	Narr		
	AM	DE-4,330,234	03-09-1995	Chwalisz		
	AN	DE-4,344,463	06-29-1995	Stockemann		
	AO	EP-022,317	01-14-1981	Watanabe		
	AP	EP-166,533	04-11-1990	Campbell		
	AQ	EP-208,510	01-14-1987	Kadin		

Examiner	Date	1
Signature	Considere	
Signature		

EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPEP 609. Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to applicant. ¹Applicant's unique citation designation number (optional). ²See Kinds Codes of USPTO Patent Documents at www.uspto.gov or MPEP 901.04. ³ Enter Office that issued the document, by the two-letter code (WIPO Standard ST.3). ⁴ For Japanese patent documents, the indication of the year of the reign of the Emperor must precede the serial number of the patent document. ⁵Kind of document by the appropriate symbols as indicated on the document under WIPO Standard ST.16 if possible. ⁶Applicant is to place a check mark here if English language Translation is attached.

This collection of information is required by 37 CFR 1.97 and 1.98. The information is required to obtain or retain a benefit by the public which is to file (and by the USPTO to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.14. This collection is estimated to take 2 hours to complete, including gathering, preparing, and submitting the completed application form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, P.O. Box 1450, Alexandria, VA 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. SEND TO: Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450.

January 29, 2004

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Filing Date

INFORMATION DISCLOSURE

First Named Inventor Zhang et al. STATEMENT BY APPLICANT Art Unit unassigned (Use as many sheets as necessary) **Examiner Name** unassigned Attorney Docket Number AHPWA1DUSA Sheet 2 of 15

			U. S. PATENT	T DOCUMENTS	
Examiner Initials*	Cite No.1	Document Number	Publication Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevant Passages or Relevant Figures Appear
-	ВІ	Number-Kind Code ² (* known) US- 5,300,655	04-05-1994	Ehrgott	
	 			von der Saal	
	BJ	US- 5,414,088	05-09-1995		
	BK	us- 5,447,928	11-05-1995	Williams	
	CA	^{US-} 5,453,516	09-26-1995	Fischer	
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	ВQ	EP-535,529	09-24-1992	Takahashi				

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Signature	Considered	
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Attorney Docket Number

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	EA	^{US-} 6,153,622	11-28-2000	Cameron			
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	cq	HU-P9600165-A	01-28-1997	Levi					

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Sheet 4 of 15

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Complete if Known					
Application Number	10/767,813				
Filing Date	January 29, 2004				
First Named Inventor	Zhang et al.				
Art Unit	unassigned				
Examiner Name	unassigned				
Attorney Docket Number	AHPWA1DUSA				

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	DM	RU-2100359	12-27-1997	Kovelman		<u> </u>
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INFORMATION DISCLOSURE STATEMENT BY APPLICANT

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Sheet 5

of 15

	Cor	inpiece ii raiowii
	Application Number	10/767,813
	Filing Date	January 29, 2004
	First Named Inventor	Zhang et al.
	Art Unit	unassigned
	Examiner Name	unassigned
1	Attorney Docket Number	AHPWA1DUSA

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	HA	^{US-} 2001-0025051-A1	09-27-2001	Cameron			
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	EL	RU-95121739-A	02-10-1998	Buzzetti		
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	EQ	WO-91/04974	04-18-1991	von der Saal		

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STATEMENT BY APPLICANT

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Examiner Initials*	Cite No.1	Document Number	Publication Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevant Passages or Relevant Figures Appear	
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		Country Code ^{3 "} Number ^{4 "} Kind Code ⁵ (if known)	MM-DD-YYYY		Or Relevant Figures Appear	T-
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	FM	WO-93/12085	06-24-1993	Boar		
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	FO	WO-94/29272	12-22-1994	Boar		
	FP	WO-95/11013	04-27-1995	Chwalisz		
	FQ	WO-95/20389	08-03-1995	Young		

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Application Number	10/767,813			
Filing Date	January 29, 2004			
First Named Inventor	Zhang et al.			
Art Unit	unassigned			
Examiner Name	unassigned			
Attorney Docket Number	AHPWA1DUSA			

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Examiner Initials*	Cite No.1	Document Number	Publication Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevant Passages or Relevant		
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Examiner Initials*	Cite No.1	Cite Foreign Patent Document	Publication Date	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevant Passages	6
		Country Code ³ Number ⁴ Kind Code ⁵ (if known)	MM-DD-YYYY		Or Relevant Figures Appear	ריי ביי
	GL	WO-95/20972	08-10-1995	Hodgen		<u> </u>
	GМ	WO-95/33746	12-14-1995	Kamireddy		
	GN	WO-96/15794	05-30-1996	Spicer		
	GO	WO-96/19458	06-27-1996	Jones		
	GP	WO-96/19997	07-04-1996	Chwalisz		
	GQ	WO-97/13767	04-17-1997	Binder		1

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Signature	Considered	
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Sheet 8

of 15

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CHDE	Filing Date	January 29, 2004	
SURE	First Named Inventor	Zhang et al.	
ICANT	Art Unit	unassigned	
y)	Examiner Name	unassigned	
	Attemory Deaket Number	ALIDIMATOLISA	

U. S. PATENT DOCUMENTS							
Examiner Initials*	Cite No.1	Document Number	Publication Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevant Passages or Relevant Figures Appear		
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Examiner Initials*	Cite No.1	Foreign Patent Document	Publication Date	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevant Passages Or Relevant Figures Appear	-6
		Country Code ³ "Number ⁴ "Kind Code ⁵ (if known)	MM-DD-YYYY			L
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-	НМ	WO-98/14436	04-09-1998	Christ		<u> </u>
	HN	WO-98/27059	06-25-1998	Kurihara		
	НО	WO-98/55116	12-10-1998	Goulet		
	HP	WO-99/10325	03-04-1999	McNutt		
	HQ	WO-99/11264	03-11-1999	Widdowson		

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Examiner	Date
Signature	Considered .

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Sheet 9

of 15

Applica	ation Number	10/767,813	
Filing C	Date	January 29, 2004	
First N	amed Inventor	Zhang et al.	
Art Uni	t	unassigned	
Examir	ner Name	unassigned	
Attorne	y Docket Number	AHPWA1DUSA	

U. S. PATENT DOCUMENTS							
Examiner Initials*	Cite No.1	Document Number	Publication Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevant Passages or Relevan Figures Appear		
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	IL.	WO-99/15500	04-01-1999	Davis		
	IM	WO-99/44608	09-10-1999	Ramachandran		_
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					Complete if Known	
Substitute for form 1449B/PTO		Application Number	10/767,813			
INF	ORMA ⁻	ΓΙΟΝ	DISCLOSURE	Filing Date	January 29, 2004	
STA	TEME	NT BY	APPLICANT	First Named Inventor	Zhang et al.	
				Group Art Unit	unassigned	
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Sheet	10	of	15	Attorney Docket Number	AHPWA1DUSA	

		NONPATENT LITERATURE DOCUMENTS	
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	AR	ANDREANI et al., "Potential antitumor agents XVII(1). Cytotoxic agents from indole derivatives and their intermediates", <i>Acta. Pharm. Nord.</i> 1990 2(6):407-414	
	AS	ARNDT et al., "New Heterocycle substituted Benzo-Fused Azine and Azole Derivatives - Useful as Selective Herbicides for Pre or Post-Emergence Application", October 10, 1988 Derwent WPI Abstract EP 311135	
	AT	BARENGOLTS et al., "Progesterone antagonist RU486 has bone-sparing effects in ovariectomized rats", <i>Bone</i> July 1995 17(1):21-25	
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	AX	CANONNE et al., "Spirocyclization of 1-(o-Aminophenyl)cycloalkanols and 1-(2'-Amino-3'-pyridinyl)cycloalkanols", <i>J. Heterocyclic Chem.</i> January-February 1989 26:113	
	AY	CHEN et al., "Synthesis and SAR of a novel series of spirobenzothlzaepine derivatives with antiprogestin activity", POI-37, 16 th Int. Cong. Het. Chem., Montana 1997	
	AZ	CHIARINO et al., "2,1-Benzisothiazoline 2,2-Dioxide and derivatives", <i>J. Heterocycl. Chem.</i> November-December 1986 23(6):1645-1649	

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STA	STATEMENT BY APPLICANT (Use as many sheets as necessary)			First Named Inventor	Zhang et al.	
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				Examiner Name	unassigned	
Sheet	11	of	15	Attorney Docket Number	AHPWA1DUSA	

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	BR	CHWALISZ et al. "Contraceptive Pack for Implantation Inhibition - Contains	
		Competitive Progesterone Antagonist and Gestagen for Sequential Oral	
		Administration.", June 29, 1995 Derwent WPI Abstract DE 4,344,463	
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	BZ	EVANS, "The steroid and thyroid hormone receptor superfamily", <i>Science</i> May 13, 1988 240(4854):889-895	

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INF	ORMAT	rion	DISCLOSURE	Filing Date	January 29, 2004	
STA	TEME	NT BY	APPLICANT	First Named Inventor	Zhang et al.	
				Group Art Unit	unassigned	
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Sheet	12	of	15	Attorney Docket Number	AHPWA1DUSA	
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	CR	FENSOME et al., "New progesterone receptor antagonists: 3,3-disubstituted-5-aryloxindoles", <i>Bio. & Med. Chem. Lett.</i> December 2, 2002 12(23):3487-3490	
	CS	FENSOME et al., "Novel 5-aryl-1,3-dihydro-indole-2-thiones: Potent, orally active progesterone receptor agonists", <i>Bio. & Med. Chem. Lett.</i> April 7, 2003 13(7):1317-1320	
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	CX	HEWAWASAM et al., "New 3-Phenyl oxindole derivatives", April 27, 2001 Abstract of Russian Patent No. 2165925	
	CY	HORWITZ et al., "Progestin, progesterone receptors, and breast cancer", Hormone Cancer, (Vedeckis ed.) Birkhaeuser: Boston, Massachusetts 1996 pp. 283-306 (abstract)	
	CZ	KEKKONEN et al., "Sequential regiment of the antiprogesterone RU486 and synthetic progestin for contraception", Fertility and Sterility October 1993 60(4):610-615	
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STA	TEME	NT BY	APPLICANT	First Named Inventor	Zhang et al.		
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	DS	KETTEL et al., "Endocrine responses to long-term administration of the	
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		September 1991 56(3):402-407	
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	DX	MAMAEV et al., "Synthesis of 4H-Thieno [3,2-B] Pyrrol-5(6H)-One" Bulletin of the	
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		Chem. March 24, 1995 60(6):1565-1582	<u> </u>
	DZ	MICHNA et al., "Differentiation therapy with progesterone antagonists", Ann. N.Y.	
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	ER	MURPHY et al., "Regression of uterine leiomyomata in response to the	
		antiprogesterone RU486", J. Clin. Endo. Metab. February 1993 76(2):513-517	

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	ES	NARR et al., "Preparation, testing, and formulation of Imidazobenzoxazinones as	
		cardiotonics", Chemical Abstracts 1988 109:22973	<u> </u>
	ET	PERLMAN et al., "20-Oxopregnacalciferols: Vitamin D Compounds that bind the	
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	EU	PFLEGEL et al., "Polarografie con 7-Chlor-5-phenly-2-thioxo-1H-2,3-dihydro-1,3,4-	
		benzotriazepinen", Pharmazie 1982 37(10):714-717	ļ. <u> </u>
	EV	SAKATA et al., "Silver halide photographic materials useful for platemaking", Chemical	
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•	EZ	TURCK et al., "On the metabolism of 3-substituted and 3,6-disubstituted pyridazines", Tetrahedron 1993 49(3):599-606	
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	FS	VERNIN et al., "Etude Dans la Serie des Radicaux Heterocycliques. Partie XV. Decomposition aprotique de 1' amino-6-ethyl-2-benzothiazole dans des substrats aromatiques et heteroaromatiques: preparation des mesityl-6- et furyl-6-ethyl-2-benzothiazoles, des sels quaternaires et des spiropyrannes correspondants", Helvetica Chimica Acta January 24, 1979 62(1/3):21-30	

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	FT	ZHI et al., "5-Aryl-1,2-Dihydrochromeno[3,4-f]quinolines: A novel class of nonsteroidal	
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